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Description

This invention relates to compositions for treating textile materials, to a method of improving the properties of textile materials, and to textile materials having improved properties. More particularly, this invention relates to aqueous dispersions which are suitable for imparting improved durable soil release properties to extile materials containing polvesters.

Textile materials containing substantial quantities of polyester fibers have been favored over all cotton garments because the polyester fibers give improved abrasion resistance and crease resistance lot the garments. However, garments containing a substantial quantity of polyester fibers have exhibited a pronounced tendency to retain soil and stains, particularly oily stains. Furthermore, these garments have been found to resist release of the soil or stains even upon exposure to repeated laundering, In all liklihood the propensity of polyester fibers or social stains are soil or stains as due to the inherently oleopholic structure of polyester fibers. In an attempt to overcome these inherent difficulties with garments made with polyester fibers, the industry has resorted to the use of a variety of textile treating compositions which provides 15 degree of soil resistance, wicking, resistance to soil redeposition and durability to repeated home laundering in aqueous systems. There remains a need in the industry, however, for a textile treating composition which provides all of the aforementioned properties and advantages and, In addition, provides better resistance to soil redeposition during fry deaning and higher fiber to fiber friction.

Chemical Abstracts Vol. 84 No. 151460k refers to the preparation of antistatic polyesters by 20 incorporating, at any stage of the polyester polymerisation, 6—15% polycoxylkylyenelglycol or its sets: derivative and 1—10 mole % (A)_x(SO_x)M(COOH)_{x-m} where A=0ZOH or (OZ)_xOH, Z is an alkylene radical, m is an integer, X is an aromatic nucleus, M is an alkali metal, and n=0, 1 or 2.

French patent specification 1803/80 describes a film- or fibre-forming copolyester derived from at least one dicarboxylic acid and at least 2 glycols wherein one of the glycols is a polylallylene coxide) and of which as a minor proportion of the structural units consists of organic radicals containing at least one sulphonate group in the form of a metal salt.

Neither of these publications describe the preparation of aqueous textile treating compositions. It has now been discovered that the need in the industry for a textile treating composition with the advantages stated above may be met by an anionic textile treating composition comprising an aqueous 30 dispersion of a polymer having a molecular weight from 2,000 to 10,000 and comprising a copolyester of ethylene glycol, polyettylene glycol having an average molecular weight of from 200 to 1000, aromatic dicarboxylic acid containing only carbon, oxygen, nydrogen and sulfur atoms. Textile metarial aromatic dicarboxylic acid containing only carbon, oxygen, nydrogen and sulfur atoms. Textile metarials containing polyester fibers which have been treated with one or more of these copolyesters exhibit sexellents oli resistance, good wicking, improved cohesion, high fiber to fiber friction, and low yard to metal friction, and excellent durability and resistance to soil redeposition during dry cleaning and home laundering.

Unless indicated otherwise, molecular weights of the polymers used in this invention are calculated by end-group analysis according to the following formula:

Molecular weight= Acid No.+Hydroxyl No.

It is essential that the polyethylene glycol used to prepare the polymers used in this invention have an average molecular weight from 200 to 1000 and preferably from 200 to 600. Polymers which do not contain groups derived from polyethylene glycol within the molecular weight range are unsuitable for imparting excellent properties to polyester textile materials treated therewith.

The polyethylene glycol may be replaced in part with monoalkylethers of polyethylene glycols, particularly the lower monoalkylethers such as methyl, ethyl, and butyl. Polypropylene glycols and higher alkylene glycols are generally not desirable because they tend to adversely effect the properties of the textile materials treated with the copolyesters. However, polyglycols which are copolymers of ethylene glycol in higher alkylene glycols may be used provided the higher alkylene glycol is only a minor portion of the polyglycol. A preferred polyethylene glycol is polyethylene glycol having a molecular weight of about 300.

The preferred aromatic dicarboxylic acid containing only carbon, hydrogen, and oxygen atoms is terepithalic acid. Illustrative examples of other dicarboxylic acids which may be used include bezzene dicarboxylic acids e.g., isophthalic acid, and phthalic acid, naphthalene dicarboxylic acids, biphenyl dicarboxylic acids, and oxydibenzoic acids.

It is also within the scope of the present invention to include a minor portion of an alighatic dicarboxylic acid such as adipic, glutaric, succinic, dimethyladipic, pimelic, azelaic, sebacic, suberic, 1,4-cyclo-haxane dicarboxylic acid, and dodecanedioic acids, in the copolymer. The specific amount of aliphatic dicarboxylic present in the copolymer is not critical, but is usually not more than about ten mole percent of the total amount of aromatic discriboxylic acid used.

Illustrative examples of sulfonated aromatic dicarboxylic acids which may be used to prepare the

copolymers employed in this invention include the alkali metal salts of 2-naphthyldicarboxy-benzene sulfonate, 1-naphthyldicarboxy-benzene sulfonate, 2-6-dimethylpheny-13-5-dicarboxy-benzene sulfonate and pheny-13-5-dicarboxy-benzene sulfonate and pheny-13-5-dicarboxy-benzene sulfonate. The preferred sulfonated salt is 5-sulfoiosphthalic acid sodium salt.

The relative amounts of (1) ethylene glycol, (2) polyethylene glycol, (3) dicarboxylic acid containing only carbon, hydrogen and oxygen, and (4) alkali metal salt of the sulfonated aromatic dicarboxylic acid used to prepare the copolymers of this invention may vary, it is preferred to use an excess of (1) and (2) over (3) and (4) so that the end-groups of the resulting copolyester are largely, but not necessity of the control of the cont

The copolyesters used in our invention may be prepared by techniques conventional in the art for the preparation of copolyesters of glycols and dicarboxylic acids.

The preferred method for preparing the copolyester comprises reacting the desired mixture of lower 20 alkyl esters (methyl, ethyl, propyl or butyl) and polyethylene glycol. The glycol esters and oligomers produced in this ester interchange reaction are then polymerised to the desired molecular weight. The ester interchange reaction may be conducted in accordance with reaction conditions generally used for ester interchange reactions. This ester interchange reaction is usually conducted at temperatures of 20°C to 21°C in the presence of an esterification catalyst. Alkanol is formed and constantly removed thus forcing the reaction to completion. The temperature of the reaction should be controlled so that ethylene glycol does not distill from the reaction mixture. Higher temperatures may be used if the reaction is conducted under pressure.

The copolyester may be prepared by polymerising a mixture of dihydroxyethyl esters of aromatic dicarboxylic acid containing only carbon, hydrogen, and oxygen atoms and an alkali metal salt of a sulfionated aromatic dicarboxylic acid containing only carbon, hydrogen, oxygen, and sulfur atoms and low molecular weight oligomers thereof, in the presence of polyethylene glycol having a molecular weight of from 200 to 1000 to form a copolyester having a molecular weight of from 200 to 1000 to form a copolyester having a molecular weight of from 2,000 to 10,000.

The catalysts used for the ester interchange reaction are those well known to the art. They include alkali and alkaline earth metals e.g. Li, Na, Ca, Mg and Transition and Group IIB metals e.g. Mn, Co, Zn and are usually added as the oxides, carbonates or acetates.

The extent of the ester interchange reaction can be followed by the amount of methanol liberated or the disappearance of the dialkyl dibasic acids in the reaction mixture as determined by high performance liquid chromatography or any other suitable technique. The ester interchange reaction should be taken to more than 90% completion. Greater than 95% completion is preferred in order to decrease the amount of sublimates obtained in the polymerization step.

Stabilizers such as phosphorous and phosphoric acid and esters thereof may be added at the end of the ester interchange step. The purpose of the stabilizer is to inhibit degradation, oxidation, and other side reactions; destroy the catalytic activity of the ester interchange catalyst; and prevent precipitation of insoluble metal carboxylates.

When the ester interchange reaction is complete, the glycol ester products are then polymerized to produce opoplymers with a molecular weight of at least 2,000. The polymerization reaction is usually conducted at temperatures from about 250°C to about 250°C in the presence of a catalyst. Higher temperatures may be used but tend to produce darker colored products. Illustrative examples of catalyst. So the subject of the produced charge to colored products. Illustrative examples of catalyst. So by hydrated antimony periodice, and ester interchange catalysts such as salts of zinc, cobalt and managants.

Excess ethylene glycol and other volatiles liberated during the reaction are removed under vacuum. The reaction is continued until the molecular weight of the polymers, as determined by end-group analysis, is at least 2,000. Higher molecular weight copolymers, for example above 3,000, are particularly desirable because they are more easily dispersed in water. Polymerization is usually stopped lust before the reaction 5m mixture becomes too viscous to agitate and before removal from the reaction vessel becomes a problem. A practical upper limit for the molecular weight is about 10,000.

The ester interchange reaction can be conducted in various ways to produce functional products. Illustrative examples of conducting the ester interchange reaction include:

(1) All of the components, dicarboxylic dialkyl esters, ethylene glycol, and polyethylene glycol, can be 60 charged at the start and ester interchanged together. This technique is illustrated in Example 1;

(2) The lower alkyl esters of the dicarboxylic acids can be charged to the mixture of ethylene glycol sequentially and ester interchanged in sequence. This technique is illustrated in Example 8. It is understood that all of the components of the copolyester must be present before the polymerization reaction occurs;

(3) The ester interchange of dimethyl terephthalate with glycol and the ester interchange of dimethyl

sulfoisophthalate, sodium salt with glycol can be conducted in separate vessels and combined for the polymerization step. Aliphatic dicarboxylic acid can be introduced in either of these separate reactions. This method is illustrated in Example 11.

The compositions of this invention are particularly useful for treating textile materials conting polyester fibers, and are particularly useful with textile materials which are 100% polyester or polyester-cotton blends. The particular form of the textile articles to be treated in accordance with the present invention is not important and includes filaments, fibers, fabrics and films.

In order to treat textile materials with the copolymers used in this invention it is essential to heat the copolymer in contact with the surface of the textile material. The continuous phase of the anionic textile to treating composition may be removed by the same or by a previous thermal treatment or it may be allowed to evaporate before thermal treatment. After the textile material is contacted with the copolyester it is heated to an elevated temperature to heat set. The temperature required to produce a durable treatment of the textile material is about 90°C or above and preferably the temperature should not exceed 50°C. Obviously the temperature should not be so high as to melt or damage the textile material being treated, so fs temperature above the melting point of the textile materials can only be applied for very short time.

It is useful, particularly when the active group or groups of the textile material or the textile treating composition are affected by atmospheric oxygen at the temperature of the thermal treatment, to carry out the thermal treatment in the presence of an antioxidantat. The antioxidant may be present as a separate compound dissolved or dispersed in the treating composition. In general, the antioxidant used may be any 20 antioxidant generally used in the art for stabilizing polyethers from thermal degradation. In considering those most suitable in the present invention it is necessary to satisfy the criteria that the antioxidant due to stable and effective at the temperature employed in the thermal treatment and that it should produce no undesirable color or odor. For example, Santonox (Trademark) Ror Irganox (Trademark) 858, pyrogallo, or zinc dietrlydithicoarbamate may be used. A combination of two or more antioxidants may give better 20 results than either antioxidant alone. Thus, for example, a mixture of zinc dinonyldithicoarbamate with 2-alpha-methylcyclohexyl-46-dimethylphenol is more effective than either antioxidant used alone.

The nature of the composition used in this invention and methods for their preparation and use will be better understood from a consideration of the following preparations and examples which are presented for illustrative purposes. All parts and percentages are by weight unless otherwise specified.

Dimethyl terephthalate 242.8 g (1,25 moles), 74.3 g (0,25 moles) dimethylsulfoisophthalate sodium salt.

Preparation 1

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6.55 g (0.0376 moles) dimethyl adipate, 374 g (6.03 moles) ethylene glycol, 0.83 g calcium acettate monohydrate, 36 g (0.12 moles) polyethylene glycol 300 and 0.019 g antimory oxide are charged to a one 3s liter 3NRB flask. The flask is equipped with an agistator, thermocouple, nitrogen inlet and a steam condenser leading to a Barrett trap extiring to a condenser, dry ice trap and hitrogen bubbler. The mixture is flushed with nitrogen and heated under a nitrogen flow of about 50—100 cc/min. The reaction mixture becomes homogeneous at 140°C and volatiles investmental per activation of the state of 1875°C and left there until the distillation stops again (15 min.). 40 The reaction mixture is then taken to 200°C and kept at this temperature for 1.5 hours. The total volatiles collected weigh 68.5 g. Phosphorous acid (0.36 g) is added and a vacuum (25 mm) is applied. The mixture is heated to 280°C over a 1 hour period. After 7 min. at 280°C the mixture becomes viscous and the product (382.6 g) is dumed into an aluminum pie pan and allowed to solidify. The product has an acid number=48.4 spanification number=444 and a hydroxyl number of 45.8.

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Preparations 2 to 6
Using the procedure described in Preparation 1 the following copolyesters are produced.

	Preparation	2	3	4	5	6		
5	DMT	292.5	292.5	242.8	1165	485.6		
	DMSI	74.3	74.3	74.3	592.5	148.6		
10	DMA	6.55	-	-	52.2	_		
	PEG-300	36.0	36.0	56.4	300	150*		
	E.G.	374	374	360.8	1482	729		
15	Ca(Ac) ₂ H ₂ O	0.63	0.63	0.63	_	1.26		
	Sb ₂ O ₃	0.019	0.019	0.019	0.152	0.038		
20	H ₃ PO ₃	0.36	0.36	0.36	2.88	0.72		
	Mn(Ac) ₂ —4H ₂ O	_	-	_	7.0	_		
	Acid number	2.4	5.3	4.6	11.0	11.8		
25	Sap number	466	468	445	416	411		
	OH number	44.7	45.6	38.1	30.0	25.1		
30	Mole wt.	2400	2200	2600	2700	3000		

DMT-Dimethyl terephthalate

DMSI-Dimethylsulfoisophthalate sodium salt

DMA-Dimethyl adipate

35 PEG-300-Polyethylene glycol (molecular weight=300)

E.G.-Ethylene glycol

Mole wt.--Molecular weight.

Preparation 7

Trappithalic acid (830.8 g), 288.2 g 5-sulfoisophthalic acid sodium salt, 21.9 g adipic acid, 399 g striyler object) 1.2.9 g adelpic acid, 399 g striyler object) 1.2.9 g adelpic acid, 399 g antirony oxide are charged to a 3NRB flash equipped with a nitrogen inlet, thermocouple, agistor, and a ban Stark trap exiting to a dry lec trap and obbler. The mixture is flushed with nitrogen and is heated to 20°C under a slight nitrogen flow. The mixture is treated at this temperature for about 27 hours. At the end of this time the reaction mixture has an acid number of five and about 490 g votatiles are recovered in the Dean Stark trap. Phosphoric acid (22.5 g 85%) is added and the mixture is heated to 27°C under a nitrogen atmosphere. After 1 hour at 27°C about 152 g votatiles are recovered and the hitrogen flow is discontinued and a vacuum (20 mm) is applied. After 10 min. under vacuum the product [1509.2 g) becomes very viscous and it is dumped into an aluminum pie pan and allowed to solidity. The product has an acid number—25, sap number—478, hydroxyl number=26.3 and %S=2.11. The molecular weight calculated from end group analysis is 2200.

55 Preparation 8

Dimethyl terephthalate 1165 g (6 moles), 738 g (11.9 moles) ethylene glycol and 7.56 g calcium acetate monohydrate are charged to a one liter 3NBB flask equipped as in the previous example. The mixture is flushed with nitrogen and heated to 200°C and kept at this temperature for a total of 13.25 hours. Volatiles (methanol) started distilling when the reaction mixture reaches 130°C. An additional 44 g ethylene glycol is added during the last 1 hour to drive the reaction to completion. A total of 362.2 g volatiles are recovered. The charge was cooled and 592.5 g (2.0 moles) dimethyl suffosiphthalate, soddium sait, 52.2 g (0.30 moles) dimethyl suffosiphthalate, soddium sait, 52.2 g (0.30 moles) dimethyl adipate, 238 g (3.85 moles) ethylene glycol, 300 g (1 mole) Carbowax 300 and 2.52 g calcium acetate monohydrate is added and reacted at temperatures up to 200°C for a period of 8.5 hours. About 10.1 g volatiles are recovered. The product is neutralized with 5.76 g phosphorous acid and 0.152 g antimony oxide is added. A vacuum (25 mm) is applied and the mixture is heated to 282°C over a 1.5 hour

^{*} Polyethylene glycol (molecular weight=600).

period during which time 464.8 g volatiles are recovered. The product 2,133 g is dumped into an aluminum pan and allowed to solidify.

Preparation 9

Dimethyl terephthalate (1165.0 g. 6 moles), 592.5 g (2 moles) dimethyl sutfoisophthalate sodium salt. 52.2 g (0.30 moles) dimethyl adipate, 300 g (1.0 moles) polyethylene glycol (mole weight 300), 1184.0 g. (19.08 moles) ethylene glycol, 3.55 g manganese acetate tetrahydrate and 0.152 g antimony trioxide are charged to a 5 liter 3 neck round bottom flask equipped with an agitator, a thermocouple, nitrogen inlet and a Dean-Stark trap exiting through a condenser, dry ice trap and a nitrogen bubbler. The charge is flushed with nitrogen and heated slowly to 200°C over a period of 10.5 hours under a slight nitrogen flow. During this time, 520 g of volatiles are collected indicating that the ester interchange reaction is about 88% completion. The mixture is cooled to 150°C and 1.44 g of phosphorus acid in 2 ml water is added. It is ne heated up to 280°C over approximately 1.5 hour period while applying a vacuum of 25 mm Hg. The product is vacuum stripped at 280°C for about 7 minutes until it became viscous. The product weighs 2106 g and to 150°C. The product is discharged into an aluminum pan and allowed to solidify. It has an acid number=10.7, hydroxyl number=21.5 and a molecular weight of \$500.

Preparation 10

Dimethyl terephthalate 1165.0 g (6 moles), 892.5 g (2 moles) dimethyl sulfoisophthalate sodium salt.

52.2 g (1.0 moles) dimethyl adipate, 300 g (1.0 moles) noveltwipen glycol (mole weight 300), 1020 g (16.4 moles) ethylene glycol, 10.08 g calcium acetate monohydrate and 0.152 g antimomy trioxide, are charged to a 5 liter 3 neck round bottom flask equipped with an agitator, thermocouple, nitrogen inter and a Dean-Stark trap exiting through a condenser, dry (se trap and a nitrogen bubbler. The charge is flushed with nitrogen and slowly heated to 200°C over a period of 12 hours. During this time, 504 g of volutiles are recovered to the control of the sets interchange reaction. The product is cooled to 150°C and 5.76 g phosphorus acid dissolved in ca. 5 ml. water is added it is then heated up to 280°C over approximately 1 hour and fifty minutes while applying a vacuum of 25 mm Hg. The product is vacuum stripped at 280°C for about 10 minutes until it becomes viscous. The product weights 2112 g and the volutiles 472 g. The product is discharged into an aluminum pan and allowed to solidify. It has an acid anumber 2.2, hydroxyl number 18.5 and molecular weight of 4300.

Preparation 11

Dimethylsulfoisophthalate, sodium salt (592.5 g. 2.00 moles), 52.2 g (0.300 moles) dimethyl adjoate, 300 g (1.00 moles) Carbowax 300, 43.7 g (7.04 moles) ethylene glycol and 2.79 g caldium acetate as monthydrate are charged to a one liter SNRB flask. The flask is equipped with an agitator, thermocouple, nitrogen inlet, and a steam condense leading to a Barrett trap exiting to a condenser, dry ice trap and nitrogen bubbler. The mixture is flushed with nitrogen and heated to 195°C under a slight nitrogen flow. It is left at this temperature for about 1.8 hours during which time about 144 g volatiles (methanol) and collected. The product is cooled to 150°C and 1.6 g phosphorus acid is added. The product is cooled and labeled 1.

Dimethyl terephthalate (1165 g, 6.00 moles), 930.8 g (15.0 moles) ethylene glycol, 3.65 g calcium acuteta monohydrate are reacted in the same manner as described above until 384.4 g volatiles are recovered (4.5 hours). The product is neutralized with 2.08 g phosphorous acid. It is cooled and tabeled II.

1 (621 g) and 866.2 g II are charged to a 3NRB flask containing an agitator, thermocouple, nitrogen inlet and takeoff to a condenser, receiver, dry les trap, vacuum gauge and vacuum pump. The mixture is flushed with nitrogen and heated to 180—200°C until it becomes homogeneous. Antimony oxide (0.1 g) is added, a vacuum (25 mn) is applied, and the mixture is heated to 280°C over a period of 52 minutes. It is left at this temperature for another five minutes until a total of 387.5 g volatile is recovered. The vacuum is broken with nitrogen and the product (1088.2 g) was dumped and allowed to solidify. It has an acid number of 7.2, saponification number of 422, and hydroxyl number of 38.5.

Preparation 12

Dimethyl terephthalate (1165 g. 6.00 moles), 592.2 g (2.00 moles) dimethylsulfoisophthalate, sodium salt, 52.2 g (0.300 moles) dimethylsulfoisophthalate, sodium salt, 52.2 g (0.300 moles) dimethylsulfoisophthalate, 300 g (1.00 moles) Carbovax 300, 1998 g (32.2 moles) ethylene glycol, 5.04 g calcium acetate monohydiate and 0.152 g antimony oxide are reacted at temperatures up to 200°C as described in experiment 10 until 506 g volatiles are recovered. Phosphorous acid, 2.88 g, is added to this product and the mixture is polymerized under vacuum to a temperature of 290°C. The product is discharged Into an aluminum pan and allowed to solidify. It has an acid number=11.2, and hydroxyl number=24.0.

Examples

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As can be seen by the data presented in the following Tables, textile materials treated with compositions of this invention have excellent physical properties. These data are obtained by the following procedure:

Anionic textile treating compositions are prepared by adding 14 parts of the indicated copolymer, 0.1

part of a bactericide, 0,16 part of antioxidant, and 0,24 part of a polyoxyethylene (9) nonylphenol as emulaifier to 85.5 parts of water at room temperature. The water is heated to 83°C (180°F) with stirring and held at that temperature until the copolymer is completely dispersed. The resulting dispersion applied to 100%, polylethaliate that test affort at the rate of 0.14 parts by weight of copolyester per 100 parts by weight of fabric. The treated fabric is cooled to 37–49°C (100–120°F), rinsed in water at 49°C (120°F), dried, and then heated to 350°F for thirty seconds to heat-set the finish. The soil release, soil redeposition, frictional and wicking properties of the treated polyester fabric are determined and shown in the following Tables III. III. Vi and V.

. Soil releas

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The purpose of this test is to measure the ability of a fabric to release stains during home laundering. (Oily stains like used motor oil, indireal oil, oilive oil, com oil, etc., are frequently employed, but many other stains such as ketchup, French dressing, mustard, chocolate syrup, grape juice, etc., may be used). Therefore the method can be used to check the performance of soil release finishes designed to facilities to the removal of soils during washing. Tests conducted before washing and again after multiple washes serve to show the durability of the finish to laundering.

Swatches of fabrics to be tested, i.e., those treated with soil release finishing agents and untreated controls, are stained with a drop of used motor oil (or other desired stain) and allowed to stand for at least five minutes so that the drop can spread out and penetrate into the fabric. Viscous or pasty stains are probed into the fabric with a spatula.

The stained fabrics are then washed in an automatic washing machine under controlled conditions using 1 gram per liter of filed (Trademark) detergent, a 95°C (120°F) was temperature, a "small load" setting and a cold water rinse. The fabrics are then dried on the "high" setting in an automatic dryer, A 2.54 cm x2.54 cm ("x-1") square section containing the stained portion of fabric is cut out and mounted. (This is the "1 wash" sample). The staining, washing and drying procedure is repeated on the remaining portion of fabric to get a "2 wash" sample. The bulk of the fabric is washed two more times, dried, then stained and washed again to prepare a "5 wash" sample. In the same way 10 or more washes may be carried out if desired.

The samples (1 wash, 2 wash, 5 wash, etc.) are then rated according to how well the stain has been go removed by a single laundering:

5=Very Good (total stain release)

4=Good (Significant but not total stain release)

3=Moderate (Moderate stain release)

2=Poor (Partial stain release)

1=Failure (No stain release)

The performance of a soil release finish is judged by comparing its rating to that of a control (unfinished) fabric. The multiple-wash samples show the degree of durability of the soil release finish to a laundarina.

TABLE II Soil release performance

45	Copolymer of Preparation No.	1	2	3	4	5	6	9	10	11	12	(A)
	1 Wash	5	5	5	5	5	5	5	5	5	5	5
50	2 Washes	5	5	5	5	5	5	4—5	5	5	5	5
	5 Washes	45	45	45	4	4-5	4—5	4	4—5	4—5	5	4
	10 Washes	4	3	2	3	2—3	3-4	4	4—5	4—5	4	1

(A) - Commercially available copolymer of ethylene glycol-polyoxyethylene glycol-terephthalic acid.

Soil redenosition

When soiled fabrics are laundered, the degree of cleanliness achieved depends upon the ability of the detergent solution used to (a) remove soils from the fibers and (b) keep those soils suspended in the solution so that they will not redeposit on the fabric.

The purpose of this test is to measure the degree of soil redeposition that occurs on a fabric during the laundering process. It can be used to test the effectiveness of fabric finishes in preventing soil redeposition.

The test is carried out by placing a 7.62×7.62 cm (3"×3") swatch of the unsoiled fabric to be tested in a 65 300 ml. Launder-Ometer (Trademark) can, adding 100 ml. of a 1 gram per liter solution of Tide (Trademark)

detergent, four (7.62×7.62 cm (3"×3") swatches of an artificially-soiled flannelette fabric, and ten 0.64 cm (14") diameter stainless sete balls for agitation. (The flanellette fabric contains a standard soil consisting of carbon black, a fatty glyceride, and mineral oil). The can is then capped and mounted in the Launder-Ometer (Trademark). When the machine is turned on the contents of the can are agitated and 5 tumbled while being heated to 80°C (176°F) and during the 45 minutes run at that temperature. During the operation soils are removed from the flannetette fabric due to the action of the detergent, and a certain amount of these soils are redeposited on the clean test fabric.

After laundering, the samples are rinsed, dried, pressed, and assessed for degree of soil redeposition according to the following rate:

- VG = Very Good (No soil pick-up)
- G =Good (slight soil pick-up)
- M =Moderate (Moderate soil pick-up)
- F =Fair (Significant soil pick-up)
 P =Poor (Excessive soil pick-up)

When soil release finishes are being evaluated, an unfinished control is included in the test for comparison. Performance is usually checked on samples of the finished fabric that have not been washed, and on samples that have been washed two and five times by a standard procedure in an automatic awasher. The performance of the washed samples gives an indication of the durability of the finish to repeated home launderings.

TABLE III Soil redeposition

25	Copolyme	er of preparation No.	12	(A)				
	0	Wash	G	F	_			
30	1	Wash	G	F				
	5	Washes	G	F				
	10	Washes	G.	F				

(A) - Commercially available copolymer of ethylene glycol-polyoxyethylene glycol-terephthalic acid.

Yarn friction

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The Atlab (Trademark) Yarn Friction Tester is used to measure frictional properties of yarns under controlled conditions. The numerical values obtained make it possible to characterize a given yarn and, to a degree, predict how that yarn will perform during certain stages of processing and after it has been woven or knitted into a fabric. The tester can be used to evaluate the performance of finishes (Unbrients, coakies) agents, etc.) simply be applying those finishes and measuring their effects on the frictional behavior of the varn.

4s Yarns to be tested are prepared, wound on cones, and conditioned for at least 16 hours in a controlled atmosphere (usually 65% relative humidity and 24°C (75°F) in the laboratory where the measurements are to be made. The test is conducted by feeding the yarn into the machine, applying a certain, predetermined tension (Ti,), and causing it to run at a selected, constant speed over a friction-producing surface. For yarn-to-wrant measurements the surface is in the form of two or three stailniess steel pins. For yarn-to-yarn so measurements a friction-producing surface is made by twisting a loop of yarn so that the moving yarn rubs against itself.

After passing over the metal or yarn surface, the tension in the yarn increases from the initial value (T₊) to a new, higher value (T₂). The increase in tension (T₂—T₁, or T₂) is directly related to the amount of friction developed at that surface. A finish applied to the yarn may increase or decrease friction compared to an 50 unfinished control yarn, and it often will improve friction uniformity to minimize tension variations in the yarn. During the test, the increase in tension, T₂ is automatically recorded on a chart.

Yarn-to-metal tests are usually run at high speeds, e.g., 10, 50, and 100 meters per minute. Average T₁ values, in grams, are read from the curves traced on the chart.

Yarn-to-yarn tests are run at much slower speeds. For example, the yarn may be moving at only one centimeter per minute. The curves produced may be smooth or they may have a stick-slip or saw-tooth pattern. Average T, values are read from the charts and reported in grams. In addition, a ± value is used to show the degree of stick-slip variation above and below the average value. (For a smooth curve the ± value would be zero). A high sick-slip value indicates a greater degree of cohesion between yarns, filaments and fibers.

In general a low yarn-to-metal friction value is desirable. Lubricants are routinely employed to reduce

friction between the yarn and guides or other metallic parts of the processing equipment. Effective lubricants permit high production speeds without damage to the yarn.

A low yarn-to-yarn friction with no stick-slip effect evident in the curve indicates that yarns, filaments or fibers will slide easily over one another. This can be desirable in a fabric since it will result in a low flex stiffness (resistance to bending) and produce a soft feel or "hand". Frequently, however, a higher yarn-to-yarn friction with some cohesion is important. In the processing of filament yarns, for example, some cohesion is needed to maintain the integrity of the yarn bundle and prevent sagging of individual filaments. Cohesion is also required to prevent slippage and permit yarns to be wound into firm, studel packages such as cones and pirns. In certain fabrics, especially delicate ones, too little cohesion can result in slippage of one yarn over another and result in serious defects.

TABLE IV

15	Copolymer of	preparation No.	12	(A)	
	Yarn to	Yarn (T _f)	72±28	12±0	
20		Metal (T _t) M/min.	44	200	
	50	M/min.	52	180	
	100	M/min.	60	180	

(A) — Commercially available copolymer of ethylene glycol-polyoxyethylene glycol-terephthalic acid.

Wicking

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The purpose of this test is to determine the vertical wicking rate of fabrics, which is a measure of their moisture transfer characteristics. It can be used to evaluate fabric finishes designed to impart hydrophilic properties to fabrics.

Strips of the fabrics to be tested (2.54×20.32 cm) (1"'×8"), all cut in the same direction of the fabric, are used in the test. If hydropholic finishes are being tested, an unfinished control fabric should be included for comparison. Unwashed fabric and fabrics that have been laundered one or more times can be included to check the durability of the finishes to washing. All fabrics to be tested are allowed to remain at ambient atmospheric conditions for at least 16 hours before testing.

The test is carried out by raising a beaker of water until the surface of the water just meets the bottom edge of the vertically-suspended fabric strip, and measuring the wicking rate by recording the diquid rise on the fabric at intervals of one, five, and ten minutes. (A suitable dye, added to the water, produces a colored solution that makes it easier to observe the exact position of the advancing liquid front. Wicking heights at a divent time are compared. The greater the wicking height the better the moisture.

transfer characteristics of the fabric. Good moisture transfer contributes to the comfort of apparel times, particularly for garments worn against the skin.

TABLE V Wicking (in cm.)

5	Copolymer of preparation No.	2	3	4	5	6	9	(A)	Un- treated
	0 Wash 1 Min.	3.0	3.5	3.5	4.5	5.5	3.5	6.0	3.0
10	5 Min.	6.0	6.0	6.0	8.0	6.0	11.0	10.0	4.0
,,,	10 Min.	8.0	7.0	8.0	9.0	8.5	13.0	11.0	5.0
	1 Wash								
15	1 Min.	8.0	8.0	8.0	9.0	9.0	9.0	10.0	7.0
	5 Min.	12.0	13.0	12.0	14.0	12.5	13.0	16.0	10.0
	10 Min.	14.0	14.0	14.0	16.0	14.0	14.0	18.0	12.0
20	2 Wash 1 Min.	5.5	6.5	6.0	8.0	7.0	7.5	9.0	5.0
	5 Min.	9.0	11.0	9.0	12.0	11.0	12.0	12.7	7.5
25	10 Min.	11.0	13.0	11.0	15.0	12.0	13.5	14.0	9.0
	5 Wash 1 Min.	9.0	8.5	9.0	10.0	9.5	9.5	10.0	8.0
30	5 Min.	12.0	12.0	13.0	14.0	12.0	13.0	14.0	11.0
	10 Min.	14.0	16.0	14.0	16.0	14.0	16.0	15.0	12.0
35	10 Wash 1 Min.	5.0	7.0	8.0	7.0	5.5	6.0	8.0	6.5
	5 Min.	7.0	10.0	11.0	9.0	9.0	9.0	11.0	9.0
40	10 Min.	9.0	12.0	12.5	11.0	11.0	11.0	13.0	9.0

(A) - Commercially available copolymer of ethylene glycol-polyoxyethylene glycol-terephthalic acid.

Having described the invention, what is desired to be secured by Patent is:

Claims

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1. An anionic textile treating composition comprising an aqueous dispersion of a copolyester having a molecular weight of from 2,000 to 10,000 and comprising a copolyester of (1) ethylene glycol, (2) polyethylene glycol having an average molecular weight of from 200 to 1000, (3) aromatic dicarboxylic acid containing only carbon, hydrogen, and oxygen atoms, and (4) an alkali metal salt of a sulfonated aromatic dicarboxylic acid containing only carbon, hydrogen, oxygen, and sulfur atoms, wherein the molar ratio of (2) to (4) is from 0.1 to 2.0, and the molar ratio of (3) to (4) is from 1 to 8.

2. A composition according to claim 1 in which the copolyester has a molecular weight of from 2,000 to 10,000, the polyethylene glycol has a molecular weight of from 200 to 600, the aromatic disenboxylic acid (3) is terephthalic acid, the alkali metal salt (4) is the sodium salt of 5-sulfoisophthalic acid, the molar ratio of (2) to (4) is from 3 to 5.

A composition according to claim 2 wherein the polyethylene glycol has a molecular weight of about
 300.

4. A composition according to any one of claims 1 to 3 wherein the copolyester is prepared by the reaction of disklyt jesters of the discraboxylic acids with a mixture of ethylene glycol and polyethylene glycol. 5. A composition according to any one of the preceding claims wherein the aromatic discraboxylic acid containing only hydrogen, carbon, and oxygen atoms is a benzene discraboxylic acid containing only hydrogen, carbon, and oxygen atoms is a benzene discraboxylic acid.

6. A composition according to claim 5 wherein the benzene dicarboxylic acid is terephthalic acid.

- 7. A composition according to any one of the preceding claims wherein the molar ratio of acid (3) to salt (4) is from 3 to 5 and the alkali metal salt is the sodium salt.
- 8. A composition according to any one of the preceding claims wherein the polyester contains an aliphatic dicarboxylic acid selected from the group consisting of glutaric, succinic, dimethyladipic, pimelic, azelaic, sebacic, suberic, 1,4-cyclohexane dicarboxylic acid, dodecanedioic and adipic acid.
 - 3. A composition according to claim 8 wherein the aliphatic dicarboxylic acid is adipic acid present in an amount of not more than 0.5 moles of adipic acid per mole of the sodium salt (4).
 - 10. A composition according to anyone of the preceding claims wherein the copolyester is prepared by an ester interchange reaction.
- 10. 11. A composition according to claim 10 wherein the copolyester is prepared by polymerizing a mixture of dihydroxyethyl esters of aromatic dicarboxylic acid containing only carbon, hydrogen, and oxygen atoms and an alkali metal salt of a sulfonated aromatic dicarboxylic acid containing only carbon, hydrogen, oxygen, and sulfur atoms and low molecular weight ofigomers thereof, in the presence polyethylene glycol having a molecular weight of from 200 to 1000 to form a copolyester having a molecular weight of from 2,000 to 10,000.
 - 12. A process of preparing a textile material which comprises contacting a textile material containing polyester with a composition according to any one of the preceding claims and heating at a temperature above 90°C.
 - 13. Textile material having in contact therewith a copolyester as defined in any one of claims 1 to 11.
 14. Textile material containing polyester having in contact therewith a copolyester as defined in any one of claims 1 to 11.

Patentansprüche

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- 1. Anionische Textilbehandlungszusammensetzung, bestehend aus einer w

 ßfrigen Dispersion eines Copolyseters mit einem Molekulargewicht von 2000 bis 10 000 und entheltend einen Copolyseter aus (1) Ethylenglycol, (2) Polyethylenglycol eines durchschnittlichen Molekulargewichts von 200 bis 1000, (3) einer aromatischen Dicarbonsäure, die nur Kohlenstoff-, Wasserstoff- und Sauerstoffstome enthält, und (4) einem Alkalimetalsalsz einer sulfonierten aromatischen Dicarbonsäure, die nur Kohlenstoff-, Wasserstoff- Sauerstoff- und Schwefelatome enthält, wobei das Molverhältnis von (3) zu (4) zwischen 0,1 und 2,0 und 38 Molverhältnis von (3) zu (4) zwischen 1 und 8 liegt- 1 und 8 lieg
- 2. Zusammensetzung nach Anspruch 1, bei welcher der Copolyester ein Molekulargewicht von 2000 bis 10 000 besitzt, das Polyethylengkoel ein Molekulargewicht von 200 bis 600 besitzt, die aromatische Dicarbonsäure (3) aus Terephthalsäure besteht, das Alkalimetallsalz (4) aus dem Netrulumsalz von 5-Suffolsophthalsäure besteht, das Molverhältnis von (2) zu (4) zwischen 0,2 und 1,0 liegt und das Molverhältnis von (3) zu (4) zwischen 3 und 5 liedt.
- Zusammensetzung nach Anspruch 2, bei welcher das Polyethylenglycol ein Molekulargewicht von ungefähr 300 aufweist.
- 4. Zusammensetzung nach einem der Ansprüche 1 bis 3, bei welcher der Copolyester hergestellt worden ist durch Umsetzung von Dialkylestern von Dicarbonsäuren mit einem Gemisch aus Ethylenglycol und Polyethylenglycol.
 5. Zusammensetzung nach einem der vorhergehenden Ansprüche, bei welcher die aromatische
- Dicarbonsäure, die nur Wasserstoff-, Kohlenstoff- und Sauerstoffatome enthält, eine Benzoldicarbonsäure ist.
- Zusammensetzung nach Anspruch 5, bei welcher die Benzoldicarbonsäure aus Terephthalsäure besteht.
- Zusammensetzung nach einem der vorhergehenden Ansprüche, bei welcher das Molverhältnis der Säure (3) zum Salz (4) zwischen 3 und 5 liegt und das Alkalimetallsalz das Natriumsalz ist.
- Zusammensetzung nach einem der vorhergehenden Ansprüche, bei welcher der Polyester eine alphafische Dicarbonsäure enthält, die ausgewählt ist aus Glutar-, Bernstein-, Dimethyladipin-, Pimelin-, Azelain-, Sebacin-, Kork-, 1.4-Cvclohexandicarbon-, Dodecandi- und Adipin- säure.
- S. Zusammensetzung nach Anspruch 8, bei welcher die aliphatische Dicarbonsäure aus Adipinsäure besteht, welche in einer Menge von nicht mehr als 0,5 Mol Adipinsäure je Mol Natriumsalz (4) vorliegt.
- besteht, welche in einer Menge von nicht mehr als 0,5 Mol Adipinsäure je Mol Natriumsalz (4) vorliegt. 10. Zusammensetzung nach einem der vorhergehenden Ansprüche, bei welcher der Copolyester durch eine Esteraustauschreaktion hergestellt worden ist.
- 11. Zusammensetzung nach Anspruch 10, bei welcher der Copolyester hergestellt worden ist durch Polymerisation eines Gemischs aus Dilvydroxyetthyelstern von anomatischen Dicarbonsäuren, die nur Kohlenstoff-, Wasserstoff- und Sauerstoffstome enthalten, und einem Alkalimetallsalz einer sulfonierten aromatischen Dicarbonsäure, die nur Kohlenstoff-, Wasserstoff-, Suacrystoff- und Schwefelatome enthält, und niedermolekularen Oligomeren davon in Gegenwart von Polyethylenglycol eines Molekulargswichts von 200 bis 1000, um einen Copolyester mit einem Molekulargswicht von 2000 bis 10 000 herzustells.
- 12. Verfahren zur Herstellung eins Textilmaterials, bei welchem ein Textilmaterial, das einen Polyester enthält; mit einer Zusammensetzung nach einem der vorhergehenden Ansprüche in Berührung gebracht und hierauf auf eine Temperatur über 90°C erhitzt wird.

- Textilmaterial, das einen mit diesem in Kontakt stehenden Copolyester nach einem der Ansprüche
 bis 11 enthält.
- 14. Polyester enthaltendes Textilmaterial, das einen mit diesem in Kontakt stehenden Copolyester nach einem der Ansprüche 1 bis 11 enthält.

Revendications

- 1. Composition anionique pour le traitement de matières textiles comprenant une dispersion aqueuse d'un copolyester ayant un poids moléculaire de 2.000 à 10.000 et comprenant un copolyester (1) d'éthylèneglycol, (2) de polyéthylèneglycol ayant un poids moléculaire de 200 à 1.000, (3) d'un acide aromatique dicarboxylique contenant uniquement des atomes de carbone, d'uydrogène et d'oxygène, et (4) d'un sel de métal alcalin d'un acide aromatique dicarboxylique sulfoné contenant uniquement des atomes de carbone, d'uydrogène, d'oxygène et de soufre, dans lequel le rapport molaire de (2) à (4) est de 1, à 2, 0 et le rapport molaire de (3) à (4) est de 1 à 8.
- 2. Composition suivant la revendication 1, dans laquelle le copolyester a un poids moléculaire de 2.000 à 10.000, le polyéthylèneglycol a un poids moléculaire de 200 à 800, l'acide aromatique dicarboxylique (3) est l'acide téréphtalique, le sel de métal alcalin (4) est le sel de sodium de l'acide 5'sulfoisophtalique, le rapport molaire de (2) à (4) est de 0,2 à 1,0, et le rapport molaire de (3) à (4) est de 3 à 5,
- 3. Composition suivant la revendication 2, dans laquelle le polyéthylèneglycol a un poids moléculaire d'environ 300.
- 4. Composition suivant l'une quelconque des revendications 1 à 3, dans laquelle le copolyester est préparé par réaction d'esters dialcoyliques des acides dicarboxyliques avec un mélange d'éthylèneglycol et de polyéth/breqlycol.
- Composition suivant l'une quelconque des revendications précédentes, dans laquelle l'acide aromatique dicarboxylique contenant uniquement des atomes d'hydrogène, de carbone et d'oxygène est un acide benzène-dicarboxylique.
 - 6. Composition suivant la revendication 5, dans laquelle l'acide benzène-dicarboxylique est l'acide téréphtalique.
- 7. Composition suivant l'une quelconque des revendications précédentes, dans laquelle le rapport molaire de l'acide (3) au sel (4) est de 3 à 5, et le sel de métal alcalin est le sel de sodium.
- 8. Composition suivant l'une quelconque des revendications précédentes, dans laquelle le polysestre contient un acide aliphatique dicarboxylique choisi dans la classe formée par l'acide glutarique, l'acide succinique, l'acide side succinique, l'acide sébacique, l'acide separatique, l'acide sébacique, l'acide subérique, l'acide 14,4-cyclohexane-dicarboxylique, l'acide dodécenacifique et l'acide adipique.
- Somposition suivant la revendication 8, dans laquelle l'acide aliphatique dicarboxylique est l'acide adipique présent en une quantité n'excédant pas 0,5 mole d'acide adipique par mole du sel de sodium (4).
 - 10. Composition suivant l'une quelconque des revendications précédentes, dans laquelle le copolyester est préparé par une réaction de transestérification.
- 11. Composition suivant la revendication 10, dans laquelle le copolyester est préparé par 40 polymérisation d'un mélange d'esters dihydroxyéthyliques d'acide aromatique dicarboxylique contenant uniquement des atomes de carbone, d'hydrogène et d'oxygène et d'un sel de métal alcalin d'un alca
 - 12. Procédé de préparation d'une matière textile, qui comprend la mise en contact d'une matière textile comprend la mise en contact d'une matière textile command un polyester avec une compositions suivant l'une quelconque des revendications précédentes et le chauffage à une température supérieure à 90°C.
- Matière textile ayant à son contact un copolyester tel que défini dans l'une quelconque des revendications 1 à 11.
 - 14. Matière textile contenant un polyester ayant à son contact un copolyester tel que défini dans l'une quelconque des revendications 1 à 11.

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